Assessment of intra-tumoral karyotypic heterogeneity by interphase cytogenetics in paraffin wax sections

S A Southern, C S Herrington

Abstract

Aim—To analyse the effect of sectioning on the assessment of karyotypic heterogeneity by interphase cytogenetics in paraffin wax embedded normal squamous epithelium and to apply the principles derived to invasive cervical carcinoma.

Methods—Normal male (n = 5) and female (n = 5) squamous epithelia were hybridised with peri-centromeric repeat probes specific for chromosomes X (DXZ1) and 17 (D17Z1) individually and in combination to assess the effect of sectioning on mono-, di-, tri-, and tetrasomic populations. Section thickness, interobserver variation and variation between different areas of the epithelium were evaluated. Invasive squamous carcinomas of the cervix (n = 5) were then hybridised with the DXZ1 probe and intratumoral heterogeneity was assessed by comparison of signal distributions obtained from different areas.

Results—The optimum section thickness for the assessment of normal epithelium was 6 μm. Variation in the expected signal number in the range 1–4 did not introduce artefactual heterogeneity at this section thickness. The sensitivity of this approach for the detection of minor subpopulations was calculated to be 13–16%, 17–18% and 10–11% for mono-, tri- and tetrasomic populations, respectively. Karyotypic heterogeneity was detected in two of the five tumours and, in one case where the populations where clustered morphologically, a minor population representing 18% was identified.

Conclusions—Interphase cytogenetic analysis of sections from paraffin wax embedded material can be used for the detection of minor subpopulations in tumours. This approach will be of particular value in the assessment of the relation between human papillomavirus infection and tumour karyotype and in the analysis of intraepithelial neoplasia.

Keywords: interphase cytogenetics, cervix, chromosome, heterogeneity.

Interphase cytogenetics involves karyotypic analysis by the demonstration of specific DNA sequences in interphase nuclei. This has several advantages over the conventional approach, which is based on the analysis of metaphase chromosomes. Cell culture is not required and hence cytological and histological material can be analysed directly. The technique is quicker and does not require cytogenetic expertise for the analysis of results as, given appropriate specificity of controls and probe characterisation, the presence of signal can be used to indicate the appropriate target sequence. This approach also has the potential for detection of subpopulations within karyotypically complex tissues and thus can be applied to the investigation of intra-tumoral heterogeneity and of mosaicism.

Numerical chromosome abnormalities can be demonstrated by in situ hybridisation with peri-centromeric repeat probes. This approach can be used to analyse individual chromosome aneusomy in tumour derived material, such as cell lines and fine needle aspirates. The application of this approach to paraffin wax embedded material is more difficult as, if whole nuclei are extracted from thick sections, morphological correlation is not possible but, if thin sections are used, morphological assessment is achieved at the expense of nuclear truncation as a result of sectioning. However, morphological correlation is a prerequisite for the analysis of intra-tumoral heterogeneity by this technique and therefore it is important to assess the information which can reliably be gained from such an approach. Although there are studies in which karyotypic heterogeneity within tumours has been demonstrated, there has been no systematic approach to the assessment of the reliability of the data produced, of the effect of sectioning in archival material or of the sensitivity of this method for the detection of minor subpopulations.

The aim of the present study was therefore to analyse the effect of sectioning on the assessment of karyotypic heterogeneity in paraffin wax embedded material. In order that data derived from the analysis of invasive tumours, which are likely to have a variably complex karyotype, could be interpreted, the effect of nuclear sectioning was modelled using normal squamous epithelium which has a stable, normal karyotype. The principles derived from this analysis were then applied to invasive cervical carcinomas.

Methods

CHOICE OF CASES

The following three groups of cases were selected from the diagnostic files of the Royal Liverpool University Hospital: (1) five normal
cervices from hysterectomy specimens from patients with no known history of cervical neoplasia; (2) five biopsy specimens of normal non-keratinising oral squamous epithelium from male patients with no history of malignant disease; and (3) five invasive cervical squamous carcinomas.

**PROBES**
The chromosome specific probes used were biotinylated D17Z1 and DXZ1 (Oncor, Gaithersburg, Maryland, USA). These probes label peri-centromeric repetitive sequences and have been localised previously to the appropriate chromosome. The DYZ3 probe (Oncor, USA) was used to confirm the presence of a Y chromosome in biopsy specimens from male patients.

**INTERPHASE CYTOGENETICS**
The method used was based on that of Hopman *et al.* Briefly, paraffin wax sections were dewaxed and pretreated for 10 minutes in 1 M NaSCN at 80°C. Sections were washed in water and then digested with 0.4% (w/v) pepsin (Sigma, Poole, Dorset, UK; P7012) in 0.2 M HCl for 20 minutes at 37°C. Following washes in water, sections were air dried and then the probe was applied at a concentration of 1 ng/ml in 2 × SSC, 10% dextran sulphate, 60% formamide, and 1 μg/ml sheared salmon sperm DNA. Denaturation was carried out at 80°C for eight minutes and hybridisation at 37°C overnight. After hybridisation, sections were washed with 60% formamide, 2 × SSC, pre-equilibrated to pH 7.0 at 42°C for 20 minutes and then phosphate buffered saline (PBS) containing 3% (w/v) bovine serum albumin (BSA) and 0.05% Tween 20 (buffer A) for 20 minutes at room temperature. Detection was performed by sequential incubation at 37°C in monoclonal anti-biotin (diluted 1 in 100), peroxidase conjugated rabbit anti-mouse (diluted 1 in 80) and peroxidase conjugated swine anti-rabbit (diluted 1 in 100) for 30 minutes each. Washes were carried out in buffer A. Signal was developed using 3,3′-diaminobenzidine (DAB)/H₂O₂. Sections were counterstained with haematoxylin and mounted in DPX.

**INTERPRETATION**
The number of signals per nucleus was counted at a total magnification of ×630 and the following rules observed: (1) all signals were counted; (2) overlapping nuclei were not counted; and (3) split signals, as defined previously, were counted as single signals. Signal number was recorded for each nucleus individually in the order of counting. Where appropriate, slides were assessed independently by two observers.

**STATISTICAL ANALYSIS**
Distributions were assessed using the Mann-Whitney U test, corrected for tied values, and the χ² test for trend. Heterogeneity was assessed by plotting dot number per nucleus against the order in which the nuclei were counted in order to retain morphological information.

**RESULTS**

**NORMAL SQUAMOUS EPITHELIUM**

**Section thickness**
The effect of section thickness was assessed by hybridising the DXZ1 probe to 4, 6 and 8 μm sections from each of the five normal cervixes. Optimum signal distributions were obtained in all cases with 6 μm and 8 μm sections (fig 1), with 58–68% of nuclei containing two signals. No difference was observed between distributions obtained by analysis of 300 nuclei from 6 μm and 8 μm sections, but 8 μm sections were less easily interpreted because of greater nuclear superimposition, particularly in the basal and para-basal layers. Therefore, 6 μm thick sections were used in the remainder of the study.

**Interobserver variation**
Two hundred nuclei from each of three separate areas from each of the five normal cervixes were counted independently by two observers. No significant difference was found between the distributions derived in each case (fig 2).

**Effect of expected signal number**
A single signal was obtained in the majority of cells with the DXZ1 probe in male but not in female epithelia. In order to model the effect of sectioning on cell populations with different karyotypes, normal male and female squamous epithelia were hybridised with DXZ1 and D17Z1, both individually and in combination. This approach produced populations with expected dot numbers of one (male epithelium, DXZ1 alone), two (female epithelium, DXZ1 alone), three (male epithelium, DXZ1 and D17Z1), and four (female epithelium, DXZ1 and D17Z1) (fig 3). Two hundred nuclei were counted from each of three separate areas from each of the five cases and frequency distributions were derived using the first 100, 150 and 200 nuclei. Hybridisation with DXZ1 alone produced a single signal in 74–81% of nuclei in male epithelium, with two signals being identified in 0–4%; 58–69% of

![Figure 1](http://mp.bmj.com/)
nuclei contained two signals and 0–2% three signals in female epithelium. Variation between cases after cohybridisation (fig 4) was related to hybridisation efficiency as there was a direct relation between the distributions obtained with the individual probes and that with the combined probe. For example, in case 1 (shown in fig 4B) 65% of nuclei contained two signals with the individual probes compared with only 58% with each probe in case 2. However, distributions derived from the three separate areas counted did not differ significantly in any case when hybridised with one or both probes. This indicates that sectioning does not introduce apparent heterogeneity artefactually in cell populations with up to four expected signals. The number of nuclei counted had no effect over the range 100–200, indicating that deviation, even in relatively small groups of nuclei, is not introduced by sectioning.

This set of experiments also permits the assessment of the sensitivity of this approach for the detection of minor subpopulations. The presence of such populations within otherwise normal epithelium was modelled by a weighted combination of the appropriate distributions from each case, followed by comparison with the disomic distribution obtained with D17Z1.

The male epithelia were used to derive sensitivities for the detection of monosomy and trisomy, and the female epithelia for tetrasomy. For example, for the evaluation of trisomy, the distribution obtained with both DXZ1 and D17Z1 in male epithelium was combined with that obtained with D17Z1 alone and the resultant distribution compared using the chi-squared test for trend with the D17Z1 distribution. This shows that a monosomic population can be detected if it represents 13–16% of the total, a trisomic population if it represents 17–18% and a tetrasomic population if it represents 10–11%.

CERVICAL CARCINOMAS
Each of five cervical carcinomas was hybridised with the DXZ1 probe alone (fig 5). In cases normal epithelium was present in the sections and gave distributions with 60–65% of nuclei containing two signals; in three cases lymphocytes within the section gave a signal distribution with 65–70% of nuclei containing two signals, indicating technical adequacy. Signals were counted from 200 nuclei in each of three previously marked areas of all five tumours by two observers. In all cases fewer than 5% of nuclei contained no signal.

Interobserver variation
No significant interobserver variation was identified in four of the cases (fig 6), but in one case there was apparent heterogeneity in one area. By plotting the signal number per nucleus as a function of the order in which the nuclei were counted, it was shown that those cells responsible for the discrepancy were clustered morphologically. Review of the slide revealed that observer 1 had counted a tumour focus that observer 2 had not, indicating that cell selection and not differences in signal interpretation was responsible for the variation: this case showed significant intra-tumoural heterogeneity (fig 7A).

Intra-tumoral heterogeneity
Significant heterogeneity between the three areas counted was statistically demonstrable in two cases and not in three (Mann-Whitney U test; fig 7). Plotting signal number as a function of counting order demonstrated the presence of discrete populations of tumour cells in one case (figs 7A and 8). The distribution obtained from area 1 is significantly different from that obtained from area 2 (p = 0.01). It can be calculated that the minor population in this case in area 1 represented approximately 18% of the nuclei counted.

Discussion
Interphase cytogenetic analysis of paraffin wax embedded material can be carried out in two ways: by extraction of whole nuclei from thick paraffin wax sections; and by using thin paraffin wax sections. The former approach allows archival material to be used but does not permit morphological correlation as tissue architecture is destroyed. Therefore, where such correlation is important, the use of thin sections is mandatory. This approach, how-
ever, leads to the production of partial nuclear profiles with consequent loss of nuclear material and hence artefactual reduction in genetic content. For interpretation of such preparations, this artefactual effect must be separated from several other variables, including variation in karyotype and in nuclear size.

In this study, the effect of sectioning was explored in two separate ways. In the first approach, section thickness was varied whilst keeping the expected signal number constant in normal squamous epithelium. The choice of section thickness is determined by the conflicting effects of nuclear sectioning and nuclear superimposition. From the data presented here, the optimum thickness in our hands is 6 μm and the proportion of nuclei containing two signals is in the range 58–69%; this is consistent with data reported by others. The use of 4 μm sections led to a significant reduction in the proportion of normal epithelial cell nuclei containing two signals; 8 μm thick sections, although giving an acceptable signal distribution, were more difficult to interpret in

Figure 3 Interphase cytogenetics of normal squamous epithelium. Hybridisation with DXZ1 alone in (A) male and (B) female epithelia gives modal signal numbers of 1 and 2, respectively. Hybridisation with both DXZ1 and D17Z1 gives variable signal numbers but nuclei containing three signals can be identified in male epithelium (C) and four signals in female epithelium (D). Note that care must be taken not to count overlapping nuclei, as shown particularly in (D).
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Figure 4  Frequency histograms illustrating the spectrum obtained from normal squamous epithelia after co-hybridisation with DXZ1 and D17Z1 in (A) male and (B) female epithelia. The two cases illustrated in each panel represent the best and worst result obtained. The distributions obtained with the combined probe reflect the hybridisation efficiency obtained using the probes individually (see Results) and demonstrate that the modal peak does not necessarily equal the expected signal number.

view of nuclear overlapping, particularly in the basal layers of the epithelium.

The second approach involved keeping the section thickness constant at 6 μm and varying the expected signal number in normal epithelium by hybridising male and female epithelia with alphoid probes specific for chromosomes X and 17, both individually and in combination. This demonstrated that there was variation between cases, including normal epithelium, but that, when different areas of the same case were analysed, there was no difference in signal distribution. These results indicate that sectioning may lead to variation in distributions between cases even when the expected signal number is the same. This is most likely the result of a combination of factors, including variation in hybridisation efficiency, differences in fixation and small variations in section thickness. In general, it was found that repeating the hybridisation gave the same distribution, indicating that variation between cases was probably because of fixation differences. This observation indicates that a particular frequency distribution cannot be regarded as indicative of the chromosome number in the intact tissue and is of importance when analysing more complex karyotypes, particularly when there is also variation in nuclear size. However, the observation that there is no variation between areas of the same case indicates that statistically significant heterogeneity is not introduced by sectioning when 100–200 nuclei are assessed from each area.

Figure 5  Interphase cytogenetics of an invasive squamous cell carcinoma (fig 7A) showing that the majority of tumour nuclei contain two or three signals.

Figure 6  Frequency histogram illustrating that no significant interobserver variation (Mann-Whitney U test) was obtained in a case with a wide range of signal numbers. This is representative of the findings in all cases analysed.
The sensitivity for detection of a trisomic population in a background of normal cells is approximately 17–18% (that is, 34–36 cells), 13–16% for a monosomic population, and 10–11% for a tetrasomic population. The sensitivities are not affected significantly by the efficiency of hybridisation, as shown by the narrow ranges. This is most likely because variation in efficiency affects both probes equally. However, sensitivity will decrease with increasing nuclear size. Therefore, the distinction between a tumour with consistent aneusomy and an aneusomic subpopulation cannot necessarily be made—for example, a tumour consisting of small cells with a trisomic subpopulation could give the same distribution as a purely trisomic tumour with large nuclei. It is therefore more appropriate simply to state that there is a statistically significant difference between normal and tumour cell populations and that this suggests the presence of an aneusomic population.

For the detection of trisomy, the sensitivity derived equates to the presence of three signals in 9–10% of cells (that is, 18–20 nuclei). This provides some objective basis for the assertion in some studies that 10–11% can be used as a cut-off point for the detection of trisomy,¹⁵ ¹⁶

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**Figure 7** Frequency histograms comparing three morphologically distinct tumour areas from all five tumours analysed. Note that there is statistically significant heterogeneity in tumours A (p < 0.001, Mann Whitney U test) and B (p < 0.001, Mann Whitney U test), but not in tumours C–E. When compared with the distributions obtained using normal epithelium (see fig 4), these data suggest that tumours B, C and E contain tetrasomic populations, and tumours A and D trisomic populations.
Figure 8 The relation between signal number and counting order in areas 1 (A) and 2 (B) from the case illustrated in fig 7B. Note that separate populations are clearly demonstrated by this approach.

but underlines the importance of not counting overlapping nuclei in order to avoid the erroneous conclusion that an aneuploid population is present in tumours where there is prominent nuclear overlap.

Application of this approach to squamous carcinomas shows that intratumoral heterogeneity is detectable if present between distinct tumour areas and that deviant populations representing as little as 18% of the total can be detected. The sensitivity of the technique to minor variations can be increased by combining morphological correlation with statistical analysis, as illustrated by one of the carcinomas analysed, and will be of particular value in the assessment of intraepithelial neoplasia. However, small foci of apparent clustering of signal number cannot be used to suggest heterogeneity as they occur frequently in sections taken from normal epithelium as a result of random clustering. In addition, the effect of variation in nuclear size must be taken into account: the larger the nucleus, the lower the signal number will be for the same karyotype. This variable can, to some extent, be controlled for morphologically and is unlikely to affect the interpretation of large subpopulations, particularly in which there has been chromosome gain. The assessment of chromosome loss is more difficult as "left shift" of the distribution obtained from a tumour by comparison with normal epithelium cannot be reliably attributed to monosomy of a proportion of the tumour cells. However, variation within tumours is less affected by this as larger tumour cells tend to contain greater numbers of signals and therefore can be interpreted reliably as heterogeneous. Probes for more than one chromosome can be used to control for variation in nuclear size as such variation affects all probes equally: any differences must therefore result from karyotypic variation and relative chromosome loss can be detected reliably.

In conclusion, therefore, paraffin wax sections can be used to demonstrate intra-tumoral heterogeneity of chromosome number. Increase in expected signal number does not alone introduce artefactual apparent heterogeneity and differences between areas of the same tumour can be interpreted as indicating heterogeneity if variation in nuclear size is taken into account. The approach described here will be particularly useful in the analysis of intraepithelial neoplasia, where morphological correlation is important.

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